A Mathematical Model for the Release, Transport, and Retention of Per- and Polyfluoroalkyl Substances (PFAS) in the Vadose Zone

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Abstract
Per- and polyfluoroalkyl substances (PFAS) are emerging contaminants of critical concern. As surfactants, PFAS tend to accumulate at air-water interfaces and may stay in the vadose zone for long times before contaminating groundwater. Yet not well understood, the extent of retention in the vadose zone has critical implications for risk management and remediation strategies. We present the first mathematical model that accounts for surfactant-induced flow and solid-phase and air-water interfacial adsorption. We apply the model to simulate PFOS (a PFAS compound of primary concern) transport in the vadose zone at a model fire-training area site impacted by aqueous film-forming foam (AFFF). Air-water interfacial adsorption is shown to have a significant impact—amplified by the low water content due to gravity drainage—total retardation factors range from 233 to 1,355 for the sand and 146 to 792 for the soil used in the study. The simulations illustrate it can take several decades or longer for PFOS to reach groundwater. Counterintuitively, the lower water content in the sand—due to stronger drainage and weaker capillary retention—leads to retardation factors greater than for the soil. Also, most PFOS is adsorbed at air-water interfaces with only 1–2% in the aqueous phase. The implications include (1) fine-texture materials could have lower retardation factors than sand due to higher retained water content, (2) soil PFAS concentrations are likely to be orders of magnitude higher than those in groundwater at source zones. Both implications are consistent with recent field observations at hundreds of AFFF-impacted sites.

1. Introduction
Per- and polyfluoroalkyl substances (PFAS)—a group of synthetic chemicals—have become emerging contaminants of critical concern. Since the late 1940s, PFAS have been used in a wide variety of products including nonstick coatings, textiles, paper products, and firefighting foams (Buck et al., 2011; Banks et al., 2013; Wang et al., 2017). Large-scale manufacturing and use of the PFAS compounds have resulted in their widespread presence in the environment. When released at the ground surface, PFAS can infiltrate into the shallow subsurface and transport through the vadose zone to contaminate groundwater (Figure 1a). Many PFAS were created as surfactants—a property that distinguishes them from traditional contaminants—when released to the environment they tend to accumulate at air-water interfaces and may stay in the vadose zone for long periods of time (Figures 1b and 1c). Except for PFAS precursors, the majority of PFAS are not known to degrade in the environment due to the strong carbon-fluorine bond (ITRC, 2018; Wang et al., 2017). While some progress on PFAS transport in groundwater has been made in the past decade, investigations in the vadose zone have only begun recently. A number of field studies have demonstrated that the vadose zone can serve as a source zone of PFAS (Anderson et al., 2019; Dauchy et al., 2019; Filipovic et al., 2015; Høisæter et al., 2019; Xiao et al., 2015; Weber et al., 2017). A major complexity of the vadose zone is the coexistence of air and water that form fluid-fluid interfaces in the soil materials. Additional complexity can be introduced when organic immiscible liquids (e.g., chlorinated solvents and hydrocarbon fuels) are present and form interfaces with air and water. The transport and fate processes of PFAS in the vadose zone in the presence of complex fluid-fluid interfaces are not well understood (SERDP, 2017). In particular, the time scale that PFAS may stay in the vadose zone due to fluid-fluid interfacial adsorption and the potential long-term release to groundwater are unknown.
The impact of fluid-fluid interfacial adsorption on PFAS retention and transport in soil was examined initially by Brusseau (2018), who employed surface tension data for PFAS including two of primary concern—perfluorooctanoic acid (PFOA) and perfluorooctanesulfonic acid (PFOS)—and measured air-water interfacial areas along with a comprehensive retention model to conduct a theoretical assessment. Additional surface-tension-based theoretical analyses of PFAS retention have since been reported (Brusseau, 2019a, 2019b; Brusseau & Van Glubt, 2019; Costanza et al., 2019; Silva et al., 2019). Miscible-displacement laboratory experiments demonstrating that adsorption of PFAS at air-water and NAPL-water interfaces can be an important retention process in soil and sand materials have also been reported (Brusseau et al., 2019; Lyu et al., 2018). A number of factors that can influence the magnitude of air-water interfacial adsorption were investigated in these studies, including water saturation, solution composition, different types of PFAS and their concentrations, and different porous media materials. To date, all of these theoretical and experimental studies employed or assumed steady-state flow conditions. The impact of transient variably saturated flow on the transport and retention of PFAS has not been considered. Yet air-water interfaces can rapidly evolve under transient variably saturated flow driven by infiltration and evapotranspiration fluxes, which can then strongly influence the transport and retention of PFAS in the vadose zone. Concomitantly—as surfactants—PFAS present in solution can modify surface tension and hence impact flow. These interactive processes lead to a system where variably saturated flow and PFAS transport and adsorption are fully coupled. Mathematical formulations and computational frameworks that represent these coupled flow and transport processes are critical for improved understanding of PFAS transport and migration in the vadose zone.

Mathematical models for the transport of surfactants have been applied in the petroleum industry since the 1970s (Pope & Nelson, 1978; UTCHEM, 2000). The application was driven by the design and evaluation of surfactant flooding technologies for enhanced oil recovery, which often requires solutions to a highly nonlinear system of three-phase flow coupled with complex phase behaviors of surfactant-oil-aqueous mixtures. Later on, surfactant-based technologies were introduced to enhance remediation of nonaqueous phase liquid (NAPLS) contaminants by solubilization or mobilization, that is, surfactant-enhanced aquifer remediation (e.g., Fountain et al., 1991; Pennell et al., 1993). Subsequently, mathematical models were developed in the environmental literature to quantify the impact of the physical and chemical processes in surfactant-enhanced aquifer remediation (Abriola et al., 1993; Delshad et al., 1996; Ji & Brusseau, 1998). Abriola et al. (1993) assumed immobile NAPLS—trapped in otherwise water-saturated pores—that dissolve into water at the NAPL-water interfaces. The transport of organic and surfactant components in a stationary configuration of NAPL and water fluid phases was simulated. Delshad et al. (1996) adopted from the surfactant-enhanced oil recovery literature the full compositional formulations with inclusion of surfactant-oil-aqueous phase behaviors. Concomitantly, models to quantify the effects of surfactants
on unsaturated water flow and solute transport in the vadose zone were also developed (Smith & Gillham, 1994) by using a modified soil characteristic curve scaled by the change of surface tension based on Leverett (1941). The formulation was shown to compare well with sand-packed column and flow cell experiments (Henry et al., 2002; Henry & Smith, 2003; Karagunduz et al., 2015; Smith & Gillham, 1999). A comprehensive review on the experimental data and numerical modeling of surfactant-induced flow in the vadose zone was provided by Henry and Smith (2003). The prior modeling studies incorporated solid-phase adsorption for the transport of surfactants, but not fluid-fluid interfacial adsorption. More recently, Costanza-Robinson and Henry (2017) modified HYDRUS-1D (Simunek et al., 2008) to simulate surfactant-induced flow to evaluate the aqueous air-water interfacial partitioning tracer test (IPTT) method. Both solid-phase and air-water interfacial adsorption were included. The adsorption coefficients—though varying among simulations—were kept constant during each simulation, and thus have underestimated the nonlinear responses of the adsorption processes under transient flow. In general, the above cited references have focused more on the impact of surfactants on fluid flow with less emphasis on the transport processes of surfactants themselves. Thus, to accurately simulate PFAS transport, there is a critical need to develop a flow and transport model that explicitly incorporates the full range of interfacial adsorption processes under transient conditions.

Compared to general surfactants, mathematical modeling studies of PFAS transport in the vadose zone are very limited. Shin et al., (2011, 2012) applied the U.S. EPA Pesticide Root Zone Model Version 3 (PRZM-3) (EPA, 2001) to simulate PFOA transport in the vadose zone. Air-water interfacial adsorption was not considered in their model. Recently, a one-dimensional (1-D) solute-transport model incorporating rate-limited solid-phase adsorption and air-water interfacial adsorption was applied to simulate the breakthrough curves of PFOS and PFOA in soil-/sand-packed column experiments under unsaturated conditions (Brusseau, 2020). These simulations focused on the transport of PFAS at steady-state unsaturated flow with a uniform saturation—no transient effects of water flow were simulated. To the best of our knowledge, no models for PFAS transport to date have accounted for both transient variably saturated flow and solid-phase and fluid-fluid interfacial adsorption processes.

Here, we develop the first mathematical model for the transport of PFAS under transient variably saturated flow in the vadose zone. Our formulation explicitly accounts for the change of surface tension by dissolved PFAS as well as PFAS adsorption at both air-water interfaces and solid grain surfaces. The variably saturated flow is modeled by the Richards equation. An advection-dispersion equation coupled with process-specific adsorption terms is formulated to model the transport of PFAS. The air-water interfacial area is parameterized as a function of water saturation based on experimental measurements. The coupled flow and transport processes are solved in a fully implicit computational framework—water pressure head and the aqueous concentration of PFAS are solved simultaneously using Newton-Raphson iterations. We apply the new model to simulate the release of PFAS to a vadose zone resulting from the use of Aqueous Film-Forming Foam (AFFF) for fire-training practices over a period of 40 years. PFOS is used as a representative PFAS because of its dominance in AFFF solutions (Houtz et al., 2013; Høisæter et al., 2019). The simulations consider two PFOS concentrations (100 mg/L vs. 1,000 mg/L), two types of porous media (sand vs. soil), and two climatic conditions (semiarid vs. humid). Rainfall and evapotranspiration data sets from Arizona and New Jersey are used as examples of semiarid and humid regions, respectively. Detailed data of hydraulic properties and air-water interfacial area for the sand and soil were measured by column experiments. Solid-phase adsorption, surface tension, and the air-water interfacial adsorption coefficients are all obtained from laboratory experiments.

The mathematical formulations of variably saturated flow and the transport processes of PFAS are presented in section 2. This is followed by a description of the numerical methods in section 3 to solve the mathematical formulations. Detailed data and parameters employed are presented in section 4. We then present the simulation results in section 5, which are followed by discussions (section 6) on the key factors that control the transport and retention of PFAS in the vadose zone. We close with concluding remarks in section 7.

### 2. Mathematical Models

We formulate a mathematical model for transient variably saturated flow coupled with the transport processes of PFAS in the vadose zone in the following two subsections (sections 2.1 and 2.2).
2.1. Variably Saturated Flow

We consider 1-D variably saturated flow in the vertical dimension of the vadose zone, which may be described by the mixed form of the Richards equation as (Pinder & Celia, 2006; Richards, 1931)

\[
\frac{\partial \theta}{\partial t} - \frac{\partial \left( K \left( \frac{\partial h}{\partial z} - 1 \right) \theta \right)}{\partial z} = 0,
\]

(1)

where \( \theta = \phi S_w \) is the volumetric water content (\(-\)). The parameter \( \phi \) is the porosity of the porous medium (\(-\)), and \( S_w \) is the water saturation (\(-\)). \( K \) is the unsaturated hydraulic conductivity (cm/s); \( h \) is the water pressure head (cm); \( z \) is the spatial coordinate (assuming positive downward; cm); and \( t \) is time (s). Here we assume that the capillary pressure head (\(-h\)) scales with surface tension \( \sigma \) following Leverett (1941) and Smith and Gillham (1994) such that

\[
h = \frac{\sigma}{\sigma_0} h_0,
\]

(2)

where \( \sigma_0 \) and \( \sigma \) are surface tensions of water without and with dissolved PFAS (dyn/cm). \( h_0 \) is the water pressure head corresponding to \( \sigma_0 \). For simplicity, the contact angle is assumed unchanged with dissolved PFAS. It is noted that our formulation can be extended to include a variable contact angle, for example, using the scalings summarized in Henry and Smith (2003).

The volumetric water content \( \theta \) is often approximated as an empirical function of \( h_0 \) (based on laboratory measurements done using water without dissolved PFAS) and referred to as soil water characteristics. Here we use the functional form proposed by van Genuchten (1980). After substituting \( h_0 = \frac{\sigma_0}{\sigma} h \), we obtain equation (3)

\[
\theta = \begin{cases} 
\theta_r + \frac{\theta_s - \theta_r}{1 + \left( \frac{\sigma - \sigma_0}{\sigma_s - \sigma_0} \right)^n}, & h < 0 \\
\theta_s, & h \geq 0
\end{cases}
\]

(3)

where \( \alpha \) (1/cm) and \( n \) (\(-\)) are fitting parameters; \( m = 1 - 1/n; \theta_r \) and \( \theta_s \) are the residual and saturated water content (\(-\)), respectively. Because surface tension \( \sigma \) is a function of the PFAS aqueous concentration, equations (1) and (3) imply that the variably saturated flow and PFAS transport are fully coupled.

The unsaturated hydraulic conductivity \( K \) can also be approximated as an empirical function (Mualem, 1976; van Genuchten, 1980) of \( h \) as

\[
K(h) = K_s S_{w,e}^{0.5} \left[ 1 - \left( 1 - S_{w,e}^{1/m} \right)^m \right]^2.
\]

(4)

where \( K_s \) is the saturated conductivity (cm/s). \( S_{w,e} = \frac{\theta - \theta_r}{\theta_s - \theta_r} \) is the effective water saturation (\(-\)).

2.2. Transport of PFAS

Transport of a PFAS or other surfactant compound in the vertical dimension of the vadose zone may be described by an advection-dispersion equation with adsorption terms (Brusseau et al., 2019; Kim et al., 1997)

\[
\frac{\partial (\theta C)}{\partial t} + \rho_b \frac{\partial C}{\partial t} + \frac{\partial C_{aw}}{\partial z} + \frac{\partial}{\partial z} \left( \theta v C \right) - \frac{\partial}{\partial z} \left( \theta D \frac{\partial C}{\partial z} \right) = 0,
\]

(5)

where \( C \) is the aqueous concentration (\( \mumol/cm^3 \)); \( C_{aw} \) is the adsorbed concentration at air-water interfaces (\( \mumol/cm^3 \)); \( C_s \) is the solid-phase adsorbed concentration (\( \mumol/g \)); \( \rho_b \) is the bulk density of the porous medium (g/cm\( ^3 \)); \( v = q/\theta \) is the interstitial pore water velocity (cm/s), where \( q = -K (\partial h/\partial z - 1) \) is the Darcy flux that will be computed from equation (1). Both \( v \) and \( q \) are positive downward. \( D = \alpha (v + \tau D_p) \) is the dispersion coefficient (cm\(^2\)/s), where \( \alpha \) is the longitudinal dispersivity (cm), \( \tau = \frac{\sigma}{\rho f c} \) is the tortuosity factor for the water phase (Millington & Quirk, 1961). \( D_p \) is the molecular diffusion coefficient in free water. We note that other partitioning processes may be relevant for some PFAS such as partitioning to the air phase and organic immiscible liquids when present (Brusseau, 2018). Here we focus on two processes—solid-phase and air-water interfacial adsorption, that are the primary processes of concern for PFOS and similar PFAS.

We first present formulations of equilibrium isotherms for \( C_s \) and \( C_{aw} \). The solid-phase adsorption \( C_s \) can be modeled by the nonlinear Freundlich isotherm as suggested by prior experimental measurements (Brusseau et al., 2019; Brusseau, 2020; Higgins & Luthy, 2006; Wei et al., 2017).
where \( K_f \) and \( N \) are fitting parameters to experimental data.

The adsorption at air-water interfaces \( C_{aw} \) has the following form

\[
C_{aw} = A_{aw} \Gamma = A_{aw} K_{aw} C,
\]

where \( A_{aw} \) is the air-water interfacial area (cm\(^2\)/cm\(^3\)). \( \Gamma = \Gamma = K_{aw} C \) is the surface excess at the air-water interface (\( \mu \text{mol/cm}^2 \)).

Air-water interfacial area comprises two general types—capillary interfaces associated with menisci between bulk air and water, and film-associated interfaces associated with wetting films surrounding grain surfaces. The combination of the two is considered the total air-water interfacial area, which is employed in the present study.

The air-water interfacial area depends on capillary pressure head, water saturation, and the imbibition and drainage history. Here, we approximate \( A_{aw} \) as a function of water saturation \( S_w \) using equation (8), which is fitted to air-water interfacial area data measured in column experiments.

\[
A_{aw} = x_2 S_w^2 + x_1 S_w + x_0,
\]

where \( x_2, x_1, \) and \( x_0 \) are fitting parameters. The parameter values for the sand and soil used in the present work are reported in section 4. Hysteresis effects are not considered.

The air-water interfacial adsorption coefficient \( K_{aw} \) can be calculated following equation (9) by relating the surface excess \( \Gamma \) to the aqueous concentration \( C \) using the Gibbs adsorption equation with the ideal dilute solution assumption (below the critical micelle concentration) (Adamson & Gast, 1997; Brusseau, 2018; Chang & Franses, 1995; Kim et al., 1997).

\[
K_{aw} = \frac{\Gamma}{C} = -\frac{1}{RTC} \frac{\partial \sigma}{\partial \ln C}.
\]

Surface tension \( \sigma \) (dyn/cm) is a function of the aqueous concentration \( C \), which can be modeled by the Szyszkowski equation (Adamson & Gast, 1997; Chang & Franses, 1995)

\[
\sigma = \sigma_0 \left[ 1 - b \ln \left( 1 + \frac{C}{a} \right) \right],
\]

where \( a \) (\( \mu \text{mol/cm}^3 \)) and \( b \) (–) are fitting parameters to experimental data. Substituting equation (10) into equation (9) yields

\[
K_{aw} = \frac{1}{RT} \frac{\sigma_0 b}{a + C}.
\]

where \( R = 8.314 \) is the universal gas constant (J/K/mol) and \( T \) is temperature (K). Equation (11) shows that \( K_{aw} \) monotonically increases as \( C \) decreases and asymptotically approaches a maximum value.

Substituting equations (6) and (7) to equation (5), we obtain an equation for the transport of a PFAS compound in the vadose zone with equilibrium adsorption processes.

\[
\frac{\partial (\theta C)}{\partial t} + \rho \frac{\partial (K_f C^N)}{\partial t} + \frac{\partial (A_{aw} K_{aw} C)}{\partial t} + \frac{\partial}{\partial z} \left( \theta D \frac{\partial C}{\partial z} \right) = 0,
\]

where \( A_{aw} \) and \( K_{aw} \) follow equations (8) and (11), respectively.

If the solid-phase and air-water interfacial adsorption are rate-limited, kinetic adsorption models need to be applied. Here we present an example formulation based on the simple two-domain sorption kinetics model (e.g., Brusseau, 1995; Cameron & Klute, 1977). More advanced models such as continuous-distribution, multirate models that incorporate a continuous distribution of domains and associated sorption/desorption rate coefficient are also available in the literature (e.g., Chen & Wagenet, 1995, 1997; Culver et al., 1997; Li & Brusseau, 2000). In the two-domain model, the solid-phase adsorption \( C_s = C_{s1} + C_{s2} \), where \( C_{s1} \)
is the adsorbed concentration (μmol/g) in the “instantaneous” sorption domain and \( C_{s,2} \) is the adsorbed concentration (μmol/g) in the kinetic sorption domain. \( C_{s,1} \) and \( C_{s,2} \) are governed by equations (13) and (14), respectively.

\[
C_{s,1} = F_s K_f C^N,
\]

\[
\frac{dC_{s,2}}{dt} = \alpha_s \left[ (1 - F_s) K_f C^N - C_{s,2} \right],
\]

where \( F_s \) is the fraction of sorbent for which sorption is instantaneous (−); \( \alpha_s \) is the first-order rate constant for kinetic solid-phase adsorption (1/s). For air-water interfacial adsorption of PFAS, diffusion from the bulk to the air-water interface has been considered as the primary controlling factor for kinetic adsorption (e.g., Brusseau, 2020; Day et al., 2007; Loppinet & Monteux, 2016; Miller et al., 2017; Sekine et al., 2004; Valkovska et al., 2004). In general, diffusive mass transfer needs to be included to model the air-water interfacial kinetic adsorption. Here we approximate using the two-domain approach by assuming that the sorption at a fraction of the air-water interfaces is instantaneous, while it is kinetic (i.e., limited by diffusive mass transfer) at the rest of the air-water interfaces. Thus, the air-water interfacial adsorption \( C_{aw} = C_{aw,1} + C_{aw,2} \), where \( C_{aw,1} \) is the adsorbed concentration at the air-water interfaces wherein the sorption is instantaneous (μmol/cm²), and \( C_{aw,2} \) is the adsorbed concentration at the air-water interfaces wherein sorption is limited by diffusive mass transfer (μmol/cm²). Under this assumption, \( C_{aw,1} \) and \( C_{aw,2} \) have the following forms

\[
C_{aw,1} = F_{aw} A_{aw} K_{aw} C,
\]

\[
\frac{dC_{aw,2}}{dt} = \alpha_{aw} \left[ (1 - F_{aw}) A_{aw} K_{aw} C - C_{aw,2} \right],
\]

where \( F_{aw} \) is the fraction of air-water interfaces for which sorption is instantaneous (−). Substituting equations (13)–(16) to equation (5) gives the transport equation for a PFAS compound with kinetic adsorption processes

\[
\frac{\partial (\theta C)}{\partial t} + \rho_b \frac{\partial (F_s K_f C^N)}{\partial z} + \rho_b \alpha_s \left[ (1 - F_s) K_f C^N - C_{s,2} \right]
+ \frac{\partial (F_{aw} A_{aw} K_{aw} C)}{\partial t} + \alpha_{aw} \left[ (1 - F_{aw}) A_{aw} K_{aw} C - C_{aw,2} \right] + \frac{\partial }{\partial z} \left( \theta D \frac{\partial C}{\partial z} \right) - \frac{\partial }{\partial z} \left( \theta E \frac{\partial C}{\partial z} \right) = 0.
\]

Equation (17) needs to be coupled with equations (14) and (16) to solve for the kinetic adsorbed concentrations \( C_{s,2} \) and \( C_{aw,2} \). In the rest of the study, we only present simulations of PFAS transport using the equilibrium adsorption models, that is, equation (12).

### 3. Numerical Methods

We apply a fully implicit numerical framework to solve the mathematical formulations in section 2, that is, equations (1) and (12), subject to initial and boundary conditions. The Richards equation (1) is solved following the mass conservative numerical scheme of Celia et al. (1990). Cell-centered finite difference and backward Euler approximations are used for the spatial and temporal discretizations. The nonlinear system of equations resulting from the discretization are solved using a Newton-Raphson iteration scheme (quadratic convergence) as opposed to the Picard iteration (linear convergence) in Celia et al. (1990). For the advection-dispersion equation (12), a first-order upwind scheme is used for the advection term and a central difference scheme is used for the diffusion term. The adsorption terms are all treated implicitly in the temporal discretization. A tolerance for the Newton updates of both equations (1) and (12) is used as a criterion to determine convergence of the nonlinear system of equations. Our numerical implementation utilizes the gridding and automatic differentiation utilities of the Matlab Reservoir Simulation Toolbox (Lie, 2019).

Flux and pressure head boundary conditions are both implemented for the top and bottom boundaries of the domain. The top boundary also considers surface evaporation and ponding conditions. Similar to the implementation in HYDRUS-1D, when the pressure head of the top numerical cell reaches below a critical value (\( h_{cr} \)), the flux condition at the top boundary switches to a fixed pressure head condition (\( h_{top} = h_{cr} \)). Ponding conditions are used when rainfall rate exceeds the rate of infiltration. When
Figure 2. Measurements and fitted curves for (a) soil water characteristics and (b) air-water interfacial area as functions of water saturation for two porous media: Accusand and Vinton soil. The markers are measurements and the lines are fitted curves. The soil water characteristics are fitted using the van Genuchten model and the air-water interfacial area are fitted using second-degree polynomials. The parameters for these curves are presented in Table 1.

4. Data and Parameters

Parameters for two types of porous media (Accusand and Vinton soil) measured by prior experiments are used in the present work. The Accusand is a commercially available natural quartz sand (UNIMIN Corp.). It has a median grain diameter of 0.35 mm and a total organic carbon content of 0.04%, Fe, Mn, and Al oxide contents of 14, 2.5, and 12 µg/g, respectively. Its clay content is negligible. The Vinton soil was collected locally in Tucson, Arizona. It consists of 54% silica, 36% feldspar, 3% amphibole, 4.7% clay minerals, and 0.1% organic carbon content. Parameters for the soil water characteristics (Figure 2a) and relative permeability for the Accusand and Vinton soil were measured by Brusseau and colleagues (unpublished). The air-water interfacial area (Figure 2b) was measured using the aqueous-IPTT method and a mass balance (aqueous) surfactant-tracer method reported in Brusseau et al. (2007), Brusseau et al. (2015), and Araujo et al. (2015). The measurements from the aqueous-IPTT and mass balance (aqueous) surfactant-tracer method are in general lower than those measured by the gas-IPTT method, especially at lower water saturations, while higher than X-ray microtomography measurements wherein roughness-impacted film-associated air-water interfaces are not accounted for (Brusseau et al., 2007). Air-water interfacial area measured by aqueous-based methods has been shown to be more representative for transport processes in the aqueous phase (Brusseau et al., 2019; Brusseau, 2019a; Lyu et al., 2018). Saturated conductivity for the Accusand and Vinton soil were measured by Brusseau and colleagues (unpublished) and from Bagour (2001), respectively. The surface tension data for PFOS solutions (Figure 3a) was reported in Brusseau (2019b). The parameter $\sigma_0$ is 71 dyn/cm, $a = 4.00 \times 10^{-3}$ µmol/cm$^3$, and $b = 0.107$. The air-water interfacial adsorption coefficient is computed via equation (11) based on the fitted Szyszkowski equation of the surface tension data. A constant temperature $T = 293.15$ K is used. The parameters for the Freundlich isotherm for the Accusand and Vinton soil were reported in Brusseau (2020) and Brusseau et al. (2019), respectively. Note that $K_f$ in Brusseau (2020) and Brusseau et al. (2019) was for PFOS aqueous concentrations in mg/L and solid-phase adsorption concentration in mg/kg. Here we have converted to the $K_f$ values for PFOS aqueous concentrations in µmol/cm$^3$ and solid-phase adsorption in µmol/g. The molecular weight of PFOS is 500.13 g/mol. The value for the longitudinal dispersivity $\alpha_L$ is estimated using the equation $\alpha_L = 0.83(\log L)^{2.414}$ proposed by Xu and Eckstein (1995), where $L$ the apparent length scale. Here we take $L$ as the length of the computational domain. The
aqueous diffusion coefficient \(D_0\) is obtained from measured values reported in Schaefer et al. (2019). The above parameters employed for the simulations in the present work are summarized in Table 1.

We consider two climatic conditions (semiarid vs. humid) to examine the impact of climate on PFAS transport in the vadose zone. Here, by “semiarid” and “humid” we focus specifically on precipitation and evapotranspiration for the climatic conditions. The site of Walnut Gulch Kendall Grasslands (Scott, 2004) in Arizona, USA, is used to represent a semiarid region, while the site of Silas Little (Clark, 2004) in New Jersey, USA, is used to represent a humid region. Precipitation and evapotranspiration data at a 30-min resolution for a period of 10 years (1 January 2005 to 31 December 2014) were downloaded from the AmeriFlux database (URL: https://ameriflux.lbl.gov) for both sites. The average annual precipitation during the 10 years are 293 mm and 1,066 mm for the two sites, respectively. The gaps (∼10%) in the data are filled using 14-day moving-average values suggested by Scott (2010). For each site, the 10-year data are repeated 3 times to generate a 40-year data set of precipitation and evapotranspiration (see Figure S2 in the SI). For simplicity, we treat all precipitation as rainfall and do not distinguish between rain and snow in our simulations. In addition, we approximate the evapotranspiration data as potential evapotranspiration to determine

**Table 1**

<table>
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<th>Parameter</th>
<th>Accusand</th>
<th>Vinton soil</th>
<th>Unit</th>
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<td>(-2,848.6)</td>
<td>(—)</td>
</tr>
<tr>
<td>(x_0)</td>
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<td>1,543.6</td>
<td>(—)</td>
</tr>
<tr>
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<td>(μmol/g)/(μmol/cm³)²N</td>
</tr>
<tr>
<td>(N)</td>
<td>0.85</td>
<td>0.81</td>
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</table>
the surface evaporation fluxes. This should be a reasonable approximation given that we do not model transpiration through plants—the surface evaporation flux should always be smaller than the measured evapotranspiration.

PFAS can be released from various types of sources including fire-training area (FTA) sites, fire response sites, industrial sites, landfills, and wastewater treatment plants (ITRC, 2018). The release from the use of AFFF at FTA sites is considered as an example source in the present work. The fire training is assumed to occur every 10 days and last for 40 years (Moody & Field, 1999, 2000). Specific information about the fire training conditions is obtained from a report by the Federal Aviation Administration (FAA, 2010) and from the literature. The report suggests two categories for the burn area, which have an average area of 385 m² and 1,318 m², respectively. For each training session, we assume that 50 to 150 gallons (189.3 to 567.8 L) of diluted AFFF aqueous solutions are applied. The volume range is consistent with values reported in Dauchy et al. (2019) at a French FTA site (400 to 800 L per training session). We note that even larger volumes (1,200 to 3,200 L) of aqueous AFFF solutions were reported elsewhere (Moody & Field, 2000). Assuming all of the AFFF solutions infiltrate into the vadose zone, it gives a total infiltration of 0.0458 cm per training session.

We use PFOS as a representative PFAS and consider two limiting concentrations of 100 and 1,000 mg/L. The range is consistent with concentrations reported in the literature. Heisæter et al. (2019) reported that 1:100 dilution of an AFFF concentrate leads to a PFOS concentration of 100 mg/L; PFOS is the dominant PFAS accounting for 88.7% of the total PFAS. Houtz et al. (2013) analyzed AFFF concentrates manufactured between 1984 and 2010 from various manufacturers; PFOS concentration was reported to range from 4.9–11.4 g/L. Assuming these AFFF concentrates are either 3% or 6%, it leads to a PFOS concentration from 147 to 684 mg/L after dilution. Each training session is assumed to last for 2 hr from 9 a.m. to 11 a.m. during that day.

5. Results
5.1. Problem Setup

We consider a 1-D domain along the vertical (z) axis. The length of the domain is 500 cm. z = 0 cm is at the land surface, and z = 500 cm is at the bottom of the domain. The groundwater table is assumed fixed at z = 482 cm. For water flow, an atmospheric boundary condition is used at z = 0 cm, namely time-dependent flux is given by the rainfall and evaporation data. The flux condition switches to a pressure head condition under surface ponding or a very dry condition (see section 3). In our simulations, the critical pressure head \( h_c \) corresponds to an effective water saturation of 0.002. The bottom boundary condition at z = 500 cm is set to a fixed pressure head \( h = 18 \) cm based on the location of the water table. A zero gradient of the aqueous PFOS concentration is used at the bottom boundary for the transport equation. In the numerical discretization, a uniform grid size is used (\( \Delta x = 0.5 \) cm). The time step size is adaptive and constrained by the maximum number of iterations and the tolerance for the Newton-Raphson iterations. Because the rainfall and evapotranspiration data have a temporal resolution of 30 min, we enforce multiple time steps to be used for each 30 min, that is, the maximum time step size is smaller than 30 min. Our simulations have converged for \( \Delta x \) and \( \Delta t \). For the nonlinear Newton-Raphson iterations, we use absolute tolerances of \( 5 \times 10^{-4} \) cm for the pressure head, \( 1 \times 10^{-3} \) for the water content \( \theta \), \( 1 \times 10^{-4} \) \( \mu \)mol/cm³ for the aqueous concentration, and 0.1 cm²/cm³ for the air-water interfacial area, all of which have to be met for convergence at every time step. The simulations are conducted in a homogeneous vadose consisting of the Accusand and the Vinton soil that are introduced in section 4. For each porous medium, we conduct eight simulations: semiarid versus humid climates, low (100 mg/L) versus high (1,000 mg/L) PFOS concentrations, and with versus without air-water interfacial adsorption.

Before simulating the transport of PFOS, we first show some illustrative simulations for variably saturated water flow only to demonstrate the spatial distribution and temporal evolution of water saturation and air-water interfacial area. We select the largest rainfall event (60 mm of total rainfall within 4 hr) from the 10-year data at the semiarid site and simulate a period of 5 days for the Accusand and a period of 60 days for the Vinton soil. The simulation starts at the same time when the largest rainfall event occurs; rainfall events following the largest one are also included if they are within the period of the simulation. The initial condition of the pressure head is set to the time-averaged pressure head over a period of 10 years computed in a separate simulation. The bottom boundary is set to a fixed pressure head at \( h = 18 \) cm. The temporal evolution of water saturation and air-water interfacial area are presented in Figure 4.
The results show that it takes ~5 days for the wetting front to reach the groundwater table in the Accusand (Figure 4a1). For the majority of the time during the 5-day simulation, the water saturation stays below 0.25 (corresponding to a water content of 0.074) and the air-water interfacial area stays above 400 cm²/cm³ (Figure 4b1) despite the fact that the conditions represent the largest rainfall event. For Vinton, the wetting front propagation is considerably slower—it takes ~60 days for the wetting front to reach the groundwater table (Figure 4a2). For the majority of the time, the water saturation is below 0.35 (corresponding to a water content of 0.138) and the air-water interfacial area is above 700 cm²/cm³ (Figure 4b2). These distributions of water saturation (or water content) and air-water interfacial area have strong implications on the transport simulations in the following two sections, namely, they amplify the impact of air-water interfacial adsorption on the retention of PFOS in the vadose zone.

5.2. Accusand

We present the simulations of PFOS release and migration in a homogeneous vadose zone consisting of a sandy media represented by the Accusand. The vertical profiles of the aqueous PFOS concentration for 10, 20, 30, 40, years are shown in Figure 5. Comparing the concentration profiles with and without interfacial adsorption (columns 1 and 3 vs. columns 2 and 4) shows that air-water interfacial adsorption greatly reduces the downward migration of PFOS. The overall time scale for PFOS to reach groundwater has increased by 1 to 2 orders of magnitude from 1 or 2 years to several decades or longer. The extent of retardation varies among different climates and PFOS concentrations. A higher PFOS concentration and a more humid climate both accelerate the migration of PFOS. Several factors are responsible for PFOS to move faster with a higher aqueous concentration. First, the air-water interfacial adsorption coefficient becomes smaller (see Figure 3). Second, the solid-phase adsorption is nonlinear with an exponent $N < 1$ and thus the effective linear adsorption coefficient will decrease for a higher aqueous concentration. Under a more humid climate, it is expected that PFOS will migrate faster because (1) more water is flowing through the vadose zone to groundwater with higher pore water velocities leading to more “leaching”, and (2) the water content is likely higher leading to less air-water interfacial adsorption. Among the four scenarios, PFOS propagates the slowest with the low PFOS concentration under a semiarid climate—the front has only migrated down 2 m after 40 years.

To show the distribution of PFOS in the different phases, we have computed the mass per bulk volume of porous medium for the aqueous phase, solid-phase adsorption, and the air-water interfacial adsorption. The results for the semiarid climate scenarios are shown in Figure 6 (the humid-climate scenarios have similar patterns and thus not shown here). The majority of the PFOS in the vadose zone is adsorbed at air-water interfaces. For example, at the maximum concentration of the plume, the air-water interfacial adsorption accounts for roughly 99% and 95% of the total PFOS in the low and high PFOS concentration scenarios, respectively. The fraction of the air-water interfacial adsorption is slightly less in the high PFOS concentration scenario because of a lower $K_{\text{air}}$. 

Figure 4. Illustrative results of the temporal evolution of water saturation and air-water interfacial area in space. Panels (a1) and (a2) are water saturations, and panels (b1) and (b2) are air-water interfacial areas. Panels (a1) and (b1) are for the Accusand, while panels (a2) and (b2) are for the Vinton soil.
We have also computed the cumulative mass of PFOS in the vadose zone. As an example, we show the high PFOS concentration case in Figure 7. The cumulative mass is normalized by the total mass of PFOS released over 40 years. Inspection of Figure 7 gives the following four observations. (1) For all cases, the total cumulative mass of PFOS increases linearly before PFOS reaches groundwater. After PFOS reaches groundwater, the cumulative mass plateaus with slight variations driven by seasonal and interannual variations of rainfall. This implies that the amount of PFOS that goes into the vadose zone is comparable to the amount that is discharged to groundwater after this point. (2) The partitioning of mass between air-water interfacial adsorption, solid-phase adsorption, and aqueous phase varies over time, primarily driven by time-dependent rainfall events. When large infiltration occurs due to rainfall events, the water saturation increases leading to a decrease in air-water interfacial area. The PFOS adsorbed at the air-water interfaces is released to the aqueous phase and the majority of that is then adsorbed to the solid surface. (3) When air-water interfacial adsorption is considered, approximately 58% and 45% of the total released PFOS remains in the vadose zone for the semiarid and humid climates, respectively. For the semiarid case, more than 93% of the mass in the vadose zone is adsorbed at the air-water interfaces. Approximately 1.5% is in the aqueous phase, and the remaining 5.5% is in the solid-phase adsorption. For the humid-climate case, ~97% is adsorbed at the air-water interfaces, 1% is in the aqueous phase, and 2% is in the solid-phase. These fractions confirm the observations made in Figure 6. (4) When air-water interfacial adsorption is not considered, the total mass left in the vadose zone is substantially lower—~7% and ~2% remain in the vadose zone for the semiarid and humid climates, respectively.
Figure 6. The four columns are the profiles of the PFOS mass per bulk volume of the porous medium for the Accusand under the semiarid climate scenario at 10, 20, 30, and 40 years, respectively. The first and second rows are simulations with a PFOS concentration of 100 and 1,000 mg/L, respectively. AW in the legend stands for air-water.

Another way to quantify the impact of the adsorption mechanisms is to use the concept of retardation factors. We compute the retardation factor $R = 1 + K_{aw}A_{aw}/\theta + \rho_b K_f C^{n-1}/\theta$ for each numerical cell within the PFOS plume for the simulations that have included air-water interfacial adsorption. A numerical cell is considered inside the plume if its aqueous PFOS concentration is greater than 0.1% of the maximum aqueous concentration in the domain. The retardation factor is then averaged in space within the plume. Finally, we compute the mean values $\bar{R}$ over 40 years for each simulation, which are reported in Table 2. The fractions of $\bar{R}$ that are associated with air-water interfacial sorption (the $K_{aw}A_{aw}/\theta$ term) and solid-phase adsorption (the $\rho_b K_f C^{n-1}/\theta$ term) are also reported; they are referred to as $f_{awia}$ and $f_{spa}$, respectively. $\bar{R}$ ranges from 233 to 1,355, and are mainly contributed by the air-water interfacial adsorption; the air-water interfacial adsorption accounts for more than 97% of the retardation. The retardation factors are consistent with the results of the plume migration in Figure 5. The large contributions of air-water interfacial adsorption are caused by the low water content and the corresponding high air-water interfacial area. To demonstrate that, we have computed the average values of the water content, air-water interfacial area, and air-water interfacial adsorption coefficient in the plume over 40 years. The water content is down to 0.037 (corresponding to a water saturation of 0.126). It is slightly lower for higher PFOS concentration because of possibly greater surfactant-induced drainage caused by a greater decrease of surface tension at a higher aqueous PFOS concentration. For example, the surface tension drops from 65.7 to 52.8 mN/m when the aqueous concentration increases from 2 to 20 mg/L. The humid climate leads to higher average water content as expected. A major difference between low and high PFOS concentrations is the value of $K_{aw}$—it decreases by 5 to 10 times from low to high PFOS concentrations, which explains the much smaller retardation factors in the high PFOS concentration scenarios.
5.3. Vinton Soil

In this section, we perform simulations in a homogeneous vadose zone consisting of a soil media represented by the Vinton soil similar to those done on the Accusand in section 5.2. The results are presented in Figures 8–10 and Table 3. They are generally similar to the Accusand results, but with three key differences. (1) The PFOS moves much slower when compared to the Accusand simulations. After 40 years, PFOS reaches groundwater for only the high PFOS concentration case under a humid climate. (2) PFOS migration is still substantially slower for the scenarios that considered the air-water interfacial adsorption compared to those with air-water interfacial adsorption turned off. However, the extent of retardation becomes smaller when compared to Accusand—the time scale to reach groundwater is only ~5 times greater when air-water interfacial adsorption is included. In addition, the scenarios without air-water interfacial adsorption also

| Table 2 |
| Retardation Factors for the Accusand for the Simulations That Have Included Air-Water Interfacial Adsorption |
| Climates | PFOS (mg/L) | \( \theta \) (—) | \( \Lambda_{aw} \) (cm²/cm³) | \( K_{aw} \) (cm³/cm²) | \( R \) (—) | \( f_{awia} \) (%) | \( f_{spa} \) (%) |
|—— | ———--- | ———— | ———— | ———— | ———— | ——— | ——— | ——— |
| Semiarid | 100 | 0.039 | 505.0 | 0.0652 | 1,355 | 99.1 | 0.9 |
| | 1,000 | 0.037 | 510.1 | 0.0069 | 233 | 97.3 | 2.7 |
| Humid | 100 | 0.048 | 472.3 | 0.0647 | 1,101 | 99.1 | 0.9 |
| | 1,000 | 0.046 | 476.3 | 0.0151 | 326 | 98.3 | 1.7 |

Note: The retardation factors are average values in the vadose zone over the entire simulation time; each \( R \) and the fractions contributed by air-water interfacial adsorption (\( f_{awia} \)) and solid-phase adsorption (\( f_{spa} \)) correspond to one simulation. The corresponding average values of water content, air-water interfacial area, and air-water interfacial adsorption coefficient are also presented.
produce much slower PFOS migration than those in the Accusand simulations. It takes ~5 years (high PFOS concentration; humid climate) to ~20 years (low PFOS concentration; semiarid climate) for the PFOS to reach groundwater. We attribute the two differences (1) and (2) to the following factors. First is that though Vinton has higher relative permeabilities because of the higher water content, its saturated hydraulic conductivity is much smaller (~18 times smaller; see Table 1). Second is that the solid-phase adsorption in Vinton is much greater (with a $K_f$ that is ~9 times larger than Accusand). The stronger solid-phase adsorption is also evident when looking at the mass fractions in the aqueous phase, solid-phase adsorption, and the air-water interfacial adsorption in Figures 9 and 10. The solid-phase adsorption accounts for about 30% and 17% of the total mass in the vadose zone for the semiarid and humid climates, respectively (Figure 10), which are much higher than the corresponding Accusand values. The fraction of the aqueous phase PFOS is still very small—1% or 2%. The remainder is adsorbed at the air-water interfaces. (3) When air-water interfacial adsorption is turned off, the cumulative PFOS in the vadose zone plateaus at a much higher value than that of the Accusand case. Approximately 60% and 20% of the total released PFOS remain in the vadose zone for the semiarid and humid climates respectively, which are almost 10 times greater than those in the Accusand scenarios.

Similar to the Accusand case, we have also computed the average retardation factors for the simulations that have included air-water interfacial adsorption to quantify the impact of the adsorption processes on the transport of PFOS (Table 3). The average values of water content, air-water interfacial area, and air-water
The retardation factors are on the order of several hundreds—though still very large—they are lower compared to the Accusand retardation factors. This is rather counterintuitive because the Vinton soil has larger interfacial areas (for the same water saturation) and also stronger solid-phase adsorption. A closer inspection reveals that the Accusand has much lower water content than the Vinton soil (more than 3 times lower), which is the main reason responsible for the Accusand’s greater retardation factors. The retardation factors decrease from low to high PFOS concentration scenarios, similar to Accusand, due to the reduction in the air-water interfacial adsorption coefficients caused by the increase in the aqueous PFOS concentration. When looking at the contributions to the retardation factors associated with the solid-phase adsorption alone ($f_{spa}$), they are ~4 times that of the Accusand $f_{spa}$ values because Vinton has a solid-phase adsorption capacity that is almost 9 times greater.

To examine the impact of surfactant-induced flow on the transport of PFOS, we have also performed simulations wherein we kept the surface tension constant (i.e., the PFOS does not impact fluid flow) and have compared the results to those that have included the feedback of PFOS to flow. The comparison shows that PFOS-induced flow has a relatively minor impact on PFOS transport and migration in the vadose zone for both Accusand and Vinton soil, especially before PFOS reaches groundwater (see Figure S3 in the SI). A closer inspection reveals that the simulated water saturation above the groundwater table is generally small, and for Accusand and Vinton soil, saturation (and thus air-water interfacial area) is not sensitive to capillary pressure head (or water pressure head) at low saturations (see Figure 2). After PFOS reaches the higher water saturation zone near the groundwater table, saturation and air-water interfacial area become more

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**Figure 9.** The four columns are the profiles of the PFOS mass per bulk volume of the porous medium for the Vinton under the semiarid climate scenario soil at 10, 20, 30, and 40 years, respectively. The first and second rows are simulations with PFOS concentration of 100 and 1,000 mg/L, respectively. AW in the legend stands for air-water.
Figure 10. Cumulative mass of PFOS in the vadose zone for the case of high PFOS concentration (1,000 mg/L) for the Vinton soil. The total mass of PFOS and the mass in the aqueous phase, air-water interfacial adsorption, and solid-phase adsorption are all presented. The first column is under a semiarid climate, while the second column is under a humid climate. The first and second rows are with and without air-water interfacial adsorption, respectively. AW in the legend stands for air-water.

sensitive to capillary pressure head. As such, the presence of PFOS surfactant—by modifying the capillary pressure head and the soil water characteristic curve—has a more noticeable impact, though still not significant, on PFOS transport. The above observation also implies that surfactant-induced flow will likely have a greater impact on PFOS transport in fine-texture porous media wherein the water saturation is higher as can be seen in Figure S3 that the impact is slightly greater for Vinton soil than that for Accusand.

Finally, we report the global mass conservation errors. For all simulations performed in the present work, the relative global mass conservation error (at the end of 40 years) is less than 0.2% for water and less than 0.005% for PFOS, respectively. These errors are small enough with no impact on PFOS transport in the vadose zone and can be further reduced if more restricted tolerances are used for the nonlinear Newton-Raphson iterations.

Table 3

<table>
<thead>
<tr>
<th>Climates</th>
<th>PFOS (mg/L)</th>
<th>(\theta)</th>
<th>(\lambda_{aw}) (cm(^2)/cm(^3))</th>
<th>(K_{aw}) (cm(^3)/cm(^2))</th>
<th>(R) (%)</th>
<th>(f_{aw}) (%)</th>
<th>(f_{spa}) (%)</th>
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</thead>
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<tr>
<td>Semiarid</td>
<td>100</td>
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<td>0.0714</td>
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<td>844.2</td>
<td>0.0264</td>
<td>153</td>
<td>90.5</td>
<td>9.5</td>
</tr>
<tr>
<td>Humid</td>
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<td>0.152</td>
<td>686.0</td>
<td>0.0699</td>
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<td>94.3</td>
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</tr>
<tr>
<td></td>
<td>1,000</td>
<td>0.150</td>
<td>690.5</td>
<td>0.0089</td>
<td>146</td>
<td>91.7</td>
<td>8.3</td>
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</tbody>
</table>

Note. The retardation factors are average values in the vadose zone over the entire simulation time; each \(R\) and the fractions contributed by air-water interfacial adsorption \((f_{aw})\) and solid-phase adsorption \((f_{spa})\) correspond to one simulation. The corresponding average values of water content, air-water interfacial area, and air-water interfacial adsorption coefficient are also presented.
6. Discussion

We have developed mathematical formulations and a fully implicit computational framework for the transport processes of PFAS under transient variably saturated flow. To the best of our knowledge, this is the first mathematical model for PFAS transport in the vadose zone that explicitly accounts for the solid-phase and air-water interfacial adsorption processes. Though we have not compared our model directly to experimental data (such data do not yet exist), the individual components of the formulations have been validated by prior experiments. The model for surfactant-induced flow has been validated by application to sand-packed column and flow cell experiments for other surfactants (Henry & Smith, 2002; Henry et al., 2002; Karagunduz et al., 2015; Smith & Gillham, 1999). The solid-phase and air-water interfacial adsorption formulations have been used to model PFAS transport in soil-/sand-packed columns under steady-state unsaturated flow. Good agreement of breakthrough curves between simulation and experiment were reported (Brusseau et al., 2019; Brusseau, 2020).

By applying the model to simulate PFAS (using PFOS as a representative PFAS) transport in a vadose zone beneath a model FTA site, we show that adsorption at air-water interfaces has a significant impact on the transport and retention of PFAS. The retardation factors—though varying with PFOS concentrations, climates, and different types of porous media—are always on the order of several hundreds to more than 1,000. We have demonstrated that such large retardation factors are mainly caused by the low water content resulting from gravity drainage. A low water content increases the retardation factor because (1) it is in the denominator of the expression for the retardation factor, and (2) it leads to a large interfacial area and thus more interfacial adsorption. We note that though our simulations used laboratory-measured parameters, the computed water content is consistent with measured water content in large lysimeters (4 m deep and 2.5 m in diameter) packed with the Vinton soil at the University of Arizona’s Karsten Center for Turfgrass Research (Carlson et al., 2003; Young et al., 1996). The water content in the lysimeters was observed to reach as low as 0.035. We expect that vadose zones with deeper groundwater tables (4.82 m set in our simulations) will have similar or even lower water contents. A shallower groundwater table may increase the overall water content if the porous medium has a thick capillary transition zone.

The other critical finding from our simulations is that the majority of the PFOS in the vadose zone is retained at the air-water interfaces—only 1% or 2% is in the aqueous phase. This implies that the total PFAS concentration in the vadose zone is likely orders of magnitude higher than that in groundwater beneath source zones. Recent field investigations have shown that measured PFAS concentrations in the soil were up to 6 or 7 orders of magnitude higher than the concentrations in the groundwater (Anderson et al., 2019; Dauchy et al., 2019). In particular, the study by Anderson et al. (2019) was based on an analysis of 324 Air Force sites across the continental United States impacted by AFFF, and higher soil PFAS concentrations were observed in the vast majority (87%) of the data collected.

Because water content and air-water interfacial areas have a major impact on the retention of PFAS transport in the vadose zone, other factors that can influence water content and the amount of air-water interfacial area will thus also have an impact on the retention. For example, a humid climate leads to a higher water content and a lower air-water interfacial area than a semi-arid climate. Another important factor is the properties of the porous medium. Our simulations show rather counterintuitively that the Vinton soil has lower PFOS retardation factors compared to the Accusand. The reason is that though the Vinton soil has a larger interfacial area than the Accusand for the same water saturation, it retains much higher water saturation than the Accusand does in the vadose zone due to stronger capillary forces. Though more comprehensive analysis for a wider range of porous media is needed to generalize the findings, our results appear to imply that a fine-texture porous medium that can retain more water in the vadose zone, for example silt or clay, could have a lower retardation factor than sand materials similar to the scenarios of the Vinton soil versus Accusand in the present work. This is supported by several prior field observations. Through the analysis of 324 Air Force sites across the continental United States impacted by AFFF, Anderson et al. (2019) showed that the ratio between the PFAS concentration in the soil and groundwater decreases with increasing clay content, which they hypothesized is due to the higher water saturation in the presence of higher clay content. In another field investigation, Dauchy et al. (2019) showed that PFAS were detected 15 m below ground despite the presence of clay layers implying that the retardation factors in clay may not be as great as what
is usually thought for other contaminants. Our modeling framework and the detailed simulations can be used to test the above hypotheses at field sites, and more generally to provide fundamental insights into the primary factors controlling the transport, migration, and retention of PFAS in the vadose zone.

Finally, we recognize that there are a few factors that are not considered in the current study that may impact PFAS transport in the vadose zone. For example, the seasonal and interannual variability of the groundwater table may influence the water saturation in the vadose zone and hence the air-water interfaces, especially when the vadose zone is relatively shallow. In that case, not including the groundwater table dynamics will likely overestimate the retention of PFAS in the vadose zone. Also, our current simulations did not consider transpiration of water through plants. For the low PFOS concentration scenarios, surface evaporation only accounts for ~16% of the total rainfall in the Accusand simulations. It is larger for Vinton, which accounts for ~44% of the total rainfall. For the high PFOS concentration scenarios, the surface evaporation is slightly lower due to more induced drainage caused by a greater reduction in surface tension. Thus, not accounting for transpiration fluxes due to plant uptake will likely overestimate the water content and thus underestimate the time scale of retention in the vadose zone. For the adsorption processes, we have presented mathematical formulations for both equilibrium and kinetic solid-phase and air-water interfacial adsorption, but we have not simulated the impact of kinetics on the transport and retention of PFAS in the vadose zone. Thus, it remains unclear whether kinetics are important under transient flow conditions, which requires further investigation. Several other factors that can add additional complexities are soil heterogeneity, multiple components of PFAS, and the transformation of PFAS precursors in the source zone. Geological heterogeneities such as macropores and fractures can generate preferential flow paths, which can accelerate the migration of PFAS in the vadose zone. Transport of multiple PFAS compounds can generate interesting interactions among the components both in terms of modifying surface tension as well as adsorption at the air-water interfaces. Though these processes are not simulated in the present work, our mathematical formulations and the computational framework can be extended to include these additional processes to examine their quantitative impact on the migration of PFAS in the vadose zone.

7. Conclusion

We have developed a novel mathematical model for the release and migration of PFAS in the vadose zone under dynamic flow conditions. The mathematical formulation accounts for transient variably saturated flow, advective and dispersive transport, and adsorption at soil grain surfaces and air-water interfaces. We have applied the model to a model source zone impacted by AFFF due to fire trainings over a period of 40 years. A series of scenarios covering a wide range of conditions are constructed, which include two porous media (sand vs. soil), two AFFF concentrations (low vs. high), and two climates (semiarid vs. humid). PFOS is used as a representative PFAS in the AFFF solutions.

The detailed simulations show that adsorption at air-water interfaces has a strong impact on the migration and retention of PFOS in the vadose zone. More importantly, air-water interfacial adsorption is amplified by the low water content (and hence large air-water interfacial areas) in the vadose zone caused by gravity drainage. The retardation factors—though varying among the different scenarios—are always on the order of several hundreds to more than 1,000. The lower water content caused by greater gravity drainage and weaker capillary retention in the sand has also led to a counterintuitive result that the retardation factors for the sand (233 to 1,355) are larger than those for the soil (146 to 792). A direct implication is that fine-texture materials such as silt or clay could have lower PFAS retardation factors compared to sand due to the higher retained water content in the vadose zone. This is supported by field evidence of decreasing soil to groundwater PFAS concentrations with increasing clay content. The large retardation factors lead to long time scales for PFOS to reach groundwater. For the conditions we simulate, it takes several decades or longer to reach a groundwater table at ~5m below the land surface. In addition, the results show that the majority of the PFOS in the vadose zone is retained at the air-water interfaces (70% or more) and the solid surfaces (5% to 30%) leaving only 1% or 2% in the aqueous phase. This implies that PFAS concentrations in the vadose zone are likely orders of magnitude higher than those in the groundwater in source zones, which is consistent with recently reported data at hundreds of AFFF-impacted FTA sites in the United States and elsewhere. Overall, the present study has provided fundamental insights into the primary factors controlling the transport and retention of PFAS in the vadose zone. More generally, the mathematical model provides a quantitative framework that can be extended to examine the complex interactions between dynamic flow and adsorption processes at field sites.
Acknowledgments
This work is in part supported by the Water, Environmental, and Energy Solutions (WEES) Initiative at the University of Arizona to B. Guo and J. Zeng, and by the NIHES Superfund Research Program (Grant P42 ES 4940) to M. L. Brusseau. We note that no new measured data were reported in this study and that all literature data used in the study have been provided in the paper except for the rainfall and evapotranspiration data, which can be downloaded from the AmeriFlux database (URL: https://ameriflux.lbl.gov).

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**Erratum**

In the originally published version, an incorrect version of Figure 3(b) displaying the parameter $K_{aw}$ was published. In Table 1, Parameters $n$ and $m$ for Vinton soil in Table 1 were inadvertently switched. The correct values are $n=3.451$ and $m=0.71$. The errors have been corrected and did not affect any of the simulation results, outcomes, or conclusions presented in the paper. This version may be considered the authoritative version of record.